Claims

1	1.	A method of modulating an immune system response to an antigen, the method
2	compr	ising steps of:
3		identifying an individual who has been or will be exposed to an antigen; and
4		administering to the individual, concurrently with exposure to the antigen, a composition
5	compr	ising at least one factor selected from the group consisting of cytokines and inducing
6	agents	, which factor is selected to bias the individual's immune response to the antigen away
7	from a	Th1 or Th2 response in a predetermined manner.
8		
9	2.	The method of claim 1, wherein:
10		the step of identifying comprises identifying an individual who is allergic to the antigen;
11	and	
12		the step of administering comprises administering a composition comprising at least one
13	factor	selected to bias the individual's immune response to the antigen away from a Th2
14	respon	ase.
15		
16	3.	The method of claim 2, wherein:
17		the step of identifying comprises identifying an individual who has previously mounted a
18	Th2 re	esponse to the antigen.
19		
20	4.	The method of claim 2, wherein:
21		the factor comprises a Th1 stimulating cytokine.

1	5.	The method of claim 2, wherein:
2		The method of claim 2, wherem. the factor is selected from the group consisting of IL-12, IL-2, IL-18, IL-1B, fragments
3	of I	L-1B, IFN α , and IFN γ .
4 5	6.	The method of claim 2, wherein: the factor comprises a Th2 stimulating cytokine.
6 7		The factor con-r
8 9 10	7.	The method of claim 2, wherein: the factor is selected from the group consisting of LPS, CD40, CD40 ligand, BCGs, ligonucleotides containing CpG motifs, TNFα, and microbial extracts.
11 12 13		8. The method of claim 7, wherein: the microbial extracts are selected from the group consisting of any Staphylococcus aureus preparation, heat killed Listeria, and modified cholera toxin.
	5 6 17	9. The method of claim 4, wherein: the step of administering comprises delivering the factor to the vicinity of T cells.
	18 19 20 21	10. The method of claim 7, wherein: the step of administering comprises delivering the factor to the vicinity of a pAPC that will internalize and display antigen to T cells.
	22	

1 2	11.	The method of claim 1, further comprising a step of: administering the antigen to the individual.
3456	12.	The method of claim 11, wherein: the step of administering the antigen comprises administering a crude antigen aration.
7 8 9	13.	the step of administering the antigen comprises administering a succession of
10 11	an	tigen.
12 13 14	14	the antigen is a polypeptide antigen; and the step of administering the antigen comprises administering a gene encoding the antigen, so that the gene becomes expressed within the individual.
	7	15. The method of claim 14, wherein: the step of administering comprises administering at least one factor that is a protein, and further comprises delivering the protein factor by administering to the individual a gene
:	20 21 22	encoding that factor. 16. The method of claim 2, wherein: Express Mail No. EJ455653983US

		ne steps of administering the antigen and administering the composition are performed			
1	the steps of administering the antigen and administering the antigen				
2	together	and comprise administering a single nucleic acid construct including genes for antigen			
3	and prot	ein factor.			
4					
5	17.	The method of claim 4, wherein:			
6		the step of administering the single nucleic acid construct comprises administering a			
7	constru	the step of administering the single state of the st			
8	single	fusion protein, containing both antigen and protein factor, is encoded.			
9					
	18.	The method of claim 2, wherein:			
10	10.	The method of claim 2, whereaster the method of claim 2, whereaster the antigen gene and the factor gene are provided on separate nucleic acid molecules.			
11					
12	10	The method of claim 2 or claim 18, wherein:			
13	19.	the antigen gene and the factor gene are coordinately regulated.			
14					
15		The method of claim 1 wherein the factor is administered in association with a targeting			
16	20.	The method of claim 1 wherein and			
17	age	nt.			
18	3	. The method of claim 11 wherein one or both of the antigen and the factor is encapsulated.			
19	9 21	The method of claim 11 wherein one of both of the			
2	0				
2	.1 22	The method of claim 11, wherein:			

together and comprise administering the antigen and composition in accounts. another. The method of claim 22, wherein: the antigen and factor are administered in association with a targeting agent. The method of claim 20 or claim 23, wherein: the targeting agent association occurs by means of an interaction selected from the group consisting of covalent bonds, hydrophobic interactions, van der Waals interactions, and				steps of administering the antigen and administering the composition are performed
another. The method of claim 22, wherein: the antigen and factor are administered in association with a targeting agent. The method of claim 20 or claim 23, wherein: the targeting agent association occurs by means of an interaction selected from the group consisting of covalent bonds, hydrophobic interactions, van der Waals interactions, and	1		the s	steps of administering the antigen and composition in association with one
The method of claim 22, wherein: the antigen and factor are administered in association with a targeting agent. The method of claim 20 or claim 23, wherein: the targeting agent association occurs by means of an interaction selected from the group consisting of covalent bonds, hydrophobic interactions, van der Waals interactions, and	2	togeth	er an	d comprise administering the antigen and composite
The method of claim 22, wherein: the antigen and factor are administered in association with a targeting agent. The method of claim 20 or claim 23, wherein: the targeting agent association occurs by means of an interaction selected from the group consisting of covalent bonds, hydrophobic interactions, van der Waals interactions, and	3	anoth	er.	
the antigen and factor are administered in association with a targeting age. The method of claim 20 or claim 23, wherein: the targeting agent association occurs by means of an interaction selected from the group consisting of covalent bonds, hydrophobic interactions, van der Waals interactions, and	4			
The method of claim 20 or claim 23, wherein: the targeting agent association occurs by means of an interaction selected from the group consisting of covalent bonds, hydrophobic interactions, van der Waals interactions, and	5	23.	Th	e method of claim 22, wherein:
The method of claim 20 or claim 23, wherein: the targeting agent association occurs by means of an interaction selected from the group consisting of covalent bonds, hydrophobic interactions, van der Waals interactions, and	6		the	e antigen and factor are administered in association
the targeting agent association occurs by means of an interaction served. the targeting agent association occurs by means of an interaction served. consisting of covalent bonds, hydrophobic interactions, van der Waals interactions, and	7			
consisting of covalent bonds, hydrophobic interactions, van der waars met	8	24.	Т	he method of claim 20 or claim 23, wherem
1: actions thereof.	9		t	ne targeting agent association occurs by means of an
1: actions thereof.	10	cor	nsistiı	ng of covalent bonds, hydrophobic interactions, van des
11 combinations as	11			ations thereof.
12	12			
13 25. The method of claim 23, wherein:	13	25	5.	The method of claim 23, wherein: the targeting agent is selected from the group consisting of mannose receptor ligand and
the targeting agent is selected from the group consists of	14			the targeting agent is selected from the group constant
the Fc receptor ligand.	15	ť	he Fc	receptor ligand.
16	16	5		
17 26. The method of claim 29, wherein:	17	7	26.	The method of claim 29, wherein:
17 26. The most of the fargeting agent comprises complement receptor ligand. 18 the targeting agent comprises complement receptor ligand.	1	8		the targeting agent comprises complement receptor figures
19	1	.9		
20 27. The method of claim 23, wherein:	2	20	27.	
the targeting agent comprises DEC205.				the targeting agent comprises DEC205.
22 Funces Mail No. EJ455653983US				2115000000

1 2	28.	,	The method of claim 23, wherein: the targeting agent comprises a ligand that interacts with a receptor on an intracellular	
3	vesi	icle	within a pAPC.	
4 5 6	29.	•	The method of claim 23, wherein: the targeting agent comprises at least the Fc portion of an Ig molecule.	
7 8 9	30).	The method of claim 23, wherein: the targeting agent comprises at least the Fc portion of an IgG molecule.	
10 11 12		31.	The method of claim 22, wherein: the step of administering comprises encapsulating the antigen and the factor together in a gle encapsulation device.	
13 14 15	i.	32	at of claim 22, wherein:	
1	7	er	ncapsulation devices.	
1	8 9 20 21		3. The method of claim 31 or 32, wherein: the step of administering the encapsulation device comprises associating the encapsulation device with a targeting agent.	
	22		DIA55653083US	

1	34.	The method of claim 33, wherein: the targeting agent is selected from the group consisting of mannose receptor ligand and
2	the Fo	receptor ligand.
3	uic i c	
4		a 1 ' 22 wherein'
5	35.	The method of claim 33, wherein:
6		the targeting agent comprises complement receptor ligand.
7		
8	36.	The method of claim 33, wherein:
	50.	the targeting agent comprises DEC205.
9		the targeting agons of 1
10		
11	37.	The method of claim 33, wherein:
12		the targeting agent directs the composition to particular vesicles within pAPCs.
13		
	20	The method of claim 33, wherein:
14	38	the targeting agent comprises at least the Fc portion of an Ig molecule.
15		the targeting agent comprises at the
16		
17	39	The method of claim 33, wherein:
18		the targeting agent comprises at least the Fc portion of an IgG molecule.
19		0. The method of claim 22, wherein:
20) 4	the step of administering comprises providing antigen and factor that are covalently
2	1	the step of administering comprises providing
2	2 1	linked to one another.
		Express Mail No. <u>EJ455653983US</u> 54

1	41.	The method of claim 22, wherein:
2		the step of administering comprises providing antigen and factor that are associated with
3	one ar	nother by means of an interaction selected from the group consisting of hydrogen bonds:
4	van d	er Waals interactions, hydrophobic interactions, and combinations thereof.
5		
6	42.	The method of claim 11, wherein:
7		the step of administering the antigen comprises administering a modified antigen.
8		
9	43.	The method of claim 42, wherein:
		the modified antigen is substantially identical to a naturally-occurring antigen that
10		tains at least one IgE binding site, but differs from that naturally-occurring antigen in that the
11	con	diffied antigen is missing at least one of the IgE binding sites.
12	mod	diffied antigen is missing at least out of
13		
14	44.	
15		the antigen comprises an autoantigen;
16		the step of identifying an individual comprises identifying an individual who has
17	mo	ounted an undesirable auto-immune response against the antigen; and
18		the factor is selected to bias the individual's immune response to the antigen away from a
19	Tì	n1 response.
20		
21	4:	The method of claim 44, wherein
22		the step of administering comprises administering a Th2 stimulating cytokine
		- National EMSS653983US

1	46.	The method of claim 44, wherein:
2		the step of administering comprises administering IL-4.
3456	47.	The method of claim 45, wherein: the step of administering comprises delivering the IL-4 to the vicinity of responding T
7 8 9	48.	The method of claim 44, wherein: the step of administering comprises administering one or more Th2 inducing agents
10 11 12	49.	The method of claim 44, wherein: the step of administering comprises administering an agent that induces IL-4 expression.
13 14	50.	A method of modulating an immune system response to an antigen, the method
15	COI	the stone of
16		isolating from an individual one or more pAPC selected from the group consisting of:
17	m	ature pAPC, immature pAPC, and precursors to pAPC; exposing the isolated cells to an antigen so that pAPC displaying the antigen are
18		exposing the isolated cells to an antigen so that processed.
19	ge	enerated, and a pre-determined set of cytokines is expressed.
20		1. The method of claim 50, further comprising:
2	1 5	1. The method of claim 50, further comp

1		administering the antigen-exposed pAPC to a subject whose immune response to the
2		is to be modulated.
3		
4	52.	The method of claim 51, wherein:
5		the antigen-exposed pAPC are mature pAPC.
6		
7	53.	The method of claim 51, wherein:
8		the antigen-exposed pAPC are immature pAPC
9		
.10	54.	The method of claim 51, wherein:
11		the pAPC are selected from the group consisting of dendritic cells, B cells, and
12	mac	crophages.
13		
14	55.	
15		the pAPC are dendritic cells.
16		
17	56	The method of claim 51, wherein:
18		the step of isolating comprises isolating immature dendritic cells from an individual; and
19		maturing the immature cells in vitro by exposure to one or more compounds selected
20	fr	om the group consisting of: GM-CSF, IL-3, and IL-4.
21		
22	2 5	7. The method of claim 53, wherein:
		Express Mail No. <u>EJ455653983US</u> April 9, 1999 57

		the step of maturing is performed concurrently with the step of exposing to antigen.
1		
2 3 4	58.	The method of claim 50, wherein: the pre-determined set of cytokines is selected from the group consisting of Th1
5	cyto	kines and Th2 cytokines.
6 7 8 9	59.	The method of claim 57, wherein: the Th1 cytokines are selected from the group consisting of IL-12, IFNα, and/or IFNγ the Th2 cytokines are selected from the group consisting of IL-4.
10 11 12	60	the step of exposing the isolated cells to an antigen comprises exposing the comprises
13	CI	rude antigen preparation.
14 15 16		The method of claim 50, wherein: the step of exposing the isolated cells to an antigen comprises exposing the cells
17	1	substantially pure antigen.
18		62. The method of claim 50, wherein:
2	.0	the antigen is a polypeptide antigen; and
2	21	the antigen is a polypeptide antigen, the the step of exposing the isolated cells to antigen comprises exposing the cells to a gene the step of exposing the isolated cells to antigen comprises exposing the cells to a gene the step of exposing the isolated cells to antigen comprises exposing the cells to a gene the step of exposing the isolated cells to antigen comprises exposing the cells to a gene the step of exposing the isolated cells to antigen comprises exposing the cells to a gene the step of exposing the isolated cells to antigen comprises exposing the cells to a gene the step of exposing the isolated cells to antigen comprises exposing the cells to a gene the step of exposing the cells to a gene the step of exposing the cells to antigen comprise exposing the cells to a gene the step of exposing the cells to antigen comprise exposing the cells to a gene the step of exposing the cells to a gene the cells to
ć.	22	encoding the antigen, so that the gene becomes expressed within the cells. Express Mail No. EJ455653983US

1 2 3	63.	the	step of exposing the cells to antigen comprises contacting the cells with an antigen ociated with a targeting agent.
4 5 6 7 8 9	iı	th a com	ne method of claim 50, wherein: ne step of exposing the isolated cells to an antigen further comprises exposing the cells position comprising a factor selected from the group consisting of cytokines and gagents, which factor is selected to bias an immune response in a subject away from a Th2 response in a pre-determined manner.
10 11 12		65.	The method of claim 64, wherein: the step of exposing comprises exposing the cells to one or more Th1 inducing agents.
13 14 15	į	66.	The method of claim 65, wherein: the Th1 inducing agents are selected from the group consisting of LPS, CD40, CD40 d, BCGs, oligonucleotides containing CpG motifs, TNFα, and microbial extracts.
1	7 8 19 20	67. aur	The method of claim 66, wherein: the microbial extracts are selected from the group consisting of any Staphylococcus preparation, heat killed Listeria, and modified cholera toxin.
	21 22	68	The method of claim 64, wherein:

1		the cytokines comprise Th1 stimulatory cytokines.
2 3 4 5	69.	The method of claim 68, wherein: the cytokines are selected from the group consisting of IL-12, IL-2, IL-18, IL-1B, tents of IL-1B, IFN α , and IFN γ .
6 7 8	70.	The method of claim 64, wherein: the step of exposing comprises exposing the cells to one or ore Th2 inducing agents.
9 10 11	71.	The method of claim 70, wherein: the Th2 inducing agents are characterized by an ability to induce IL-4 expression in the
12	pA	PC.
13 14 15	72	The method of claim 64, wherein: the cytokines comprise Th2 stimulatory cytokines.
16 17 18		The method of claim 64, wherein: the cytokines comprise IL-4.
19 20 2	o 7	74. The method of claim 64, wherein: the factor is a polypeptide; and

		the step of exposing the cells to a composition comprising the factor comprises			
1		the step of exposing the constraints the factor			
2	contac	ting the cells with a gene encoding the factor.			
3					
4	75 .	The method of claim 74, wherein:			
5		The method of Claim 7 is the series of the gene encoding the factor are coordinately the gene encoding the antigen and the gene encoding the factor are coordinately			
6 .	regul	ated.			
7					
8	76.	The method of claim 74, wherein:			
9		The method of claim 74, wherein the method of claim 74, which is the method of claim 74, which is the method of claim 74, which is the method of claim 74, wherein the method of claim 74, which is the			
10	nuc	leic acid molecule.			
11					
12	. 77.	The method of claim 76, wherein:			
13		The method of claim 76, wherein the gene encoding the factor are linked together so that the gene encoding the antigen and the gene encoding the factor are linked together so that			
14	a f	fusion protein is encoded.			
15					
16	78	The method of claim 74, wherein:			
17		The method of claim 74, wherems the gene encoding the antigen and the gene encoding the factor are provided on separate			
18	; n	nucleic acid molecules.			
19)				
2	0	79. The method of claim 64, wherein:			
	.1	79. The method of claims 5.7, the one or both of the antigen and factor are associated with a targeting agent.			
	22				
_		71455652083115			

1 2	80.		ne method of claim 79, wherein: ne association with the targeting agent occurs by means of an interaction selected from no consisting of covalent bonds, hydrogen bonds, van der Waals interactions,
3	the	grou	consisting of covalent bonds, hydrogen consisting the covalent bonds and hydrogen consisting the covalent bonds are consistent bonds.
4	hy	droph	obic interactions, and combinations thereof.
5			
6	8	۱.	The method of claim 79, wherein:
7			The method of claim 79, wherein. the targeting agent is selected from the group consisting of mannose receptor ligand and
8	ti	he Fc	receptor ligand.
9 10 11	į	82.	The method of claim 79, wherein: the targeting agent comprises complement receptor ligand.
12 13 14		83.	The method of claim 79, wherein: the targeting agent comprises DEC205.
15 16 17	5	84.	The method of claim 79, wherein: the targeting agent is capable of targeting to intracellular vesicles within pAPCs.
1	8 9 20	85	The method of claim 79, wherein: the targeting agent comprises at least the Fc portion of an Ig molecule.
	21	8	6. The method of claim 79, wherein: Express Mail No. EJ455653983US
			10.1000

1		the targeting agent comprises at least the Fc portion of an IgG molecule.
2 3 4	87.	The method of claim 50, wherein: the antigen is encapsulated.
5 6 7 8	88.	The method of claim 64, wherein: the step of exposing comprises providing the antigen and factor together in an psulation device.
9 10 11 12	89.	The method of claim 64, wherein: the step of administering comprises providing the antigen and the factor in separate apsulation devices.
13 14 15 16	90 as	The method of claim 87, 88, or 89, wherein: the step of exposing comprises exposing the cells to the encapsulation device in speciation with a targeting agent.
17 18 19 20		The method of claim 90, wherein: the targeting agent is selected from the group consisting of mannose receptor ligand and the Fc receptor ligand.
21 22		92. The method of claim 90, wherein: Express Mail No. EJ455653983US

1		the ta	rgeting agent comprises complement reco	eptor ligand.
2				
3	93.		method of claim 90, wherein:	
4		the t	targeting agent comprises DEC205.	
5 6 7	94.	The	e method of claim 90, wherein: targeting agent is capable of targeting to	particular vesicles within pAPCs.
8 9 10	95.	Th	ne method of claim 90, wherein: ne targeting agent comprises at least the Fo	portion of an Ig molecule.
11 12 13	96	j. T	The method of claim 90, wherein: he targeting agent comprises at least the I	c portion of an IgG molecule.
14 15 16				antigen and factor that are associated with one the group consisting of: covalent bonds,
17 18	8	anothe hydrog	r by means of an interaction servingen bonds, van der Waals interactions, hy	drophobic interactions, and combinations thereof.
	9	98.	The method of claim 50, wherein:	the calls to a modified antigen.
	21		the step of exposing the antigen compr	ses exposing the cells to a modified antigen.
:	22			Express Mail No. <u>EJ455653983US</u>

1 2 3	99.	The method of claim 64, wherein: the antigen comprises an autoantigen; the factor is selected to bias the immune response to the antigen away from a Th1
4	respo	nse.
5		
6	100.	The method of claim 99, wherein:
7		the factor comprises a Th2 inducing agent.
8		
9	101	The method of claim 99, wherein
	•	the factor comprises an agent that induces IL-4 expression in the pAPC.
10		(IIC lactor of 1
11		c4 -lamoin'
12	10	
13		the antigen comprises an allergen; and
		the antigen comprises an array of the antigen away from a Th2 the factor is selected to bias the immune response to the antigen away from a Th2
14		
15	re	sponse.
16		a de la companya de
17	7 1	03. The method of claim 102, wherein:
13	8	the factor comprises a Th1 inducing agent.
1	9	
2	20	104. The method of claim 102, wherein:
		the factor is selected from the group consisting of LPS, CD40, CD40 ligand, D000,
	21	oligonucleotides containing CpG motifs, TNFα, and microbial extracts.
ļ	22	express Mail No. <u>EJ455653983US</u>
		Express Man 1-1

1 2 3	105.	the	e method of claim 104, wherein: microbial extracts are selected from the group consisting of any Staphylococcus sparation, heat killed Listeria, and modified cholera toxin.
4			
5	106.	Th	ne method of claim 51, wherein:
6		th	ne method of claim 31, wherem the method of claim 31, wherem the step of administering further comprises administering a cytokine selected from the
7	gro	up co	ne step of administering further compared to the subject.
8			
9	10'	7 . 7	The method of claim 106, wherein:
		1	The method of claim 106, wherem the Th1 stimulatory cytokines are selected from the group consisting of IL-12, IL-2, IL- the Th1 stimulatory cytokines are selected
10 11	18	8, IL-1	the Th1 stimulatory cytokines are serected in the Th2 stimulatory cytokines are selected in the IB, fragments of IL-1B, IFNα, and IFNγ and the Th2 stimulatory cytokines are selected
12			ne group consisting of IL-4.
13		- 4	The method of claim 51 or claim 101, further comprising:
14	1	.80	
15			administering antigen to the subject.
16			response to an antigen, the method
17		109.	A method of modulating an immune system response to an antigen, the method
18	3	comp	prising steps of: isolating from an individual one or more APC selected from the group consisting of:
19	9		isolating from an individual one of mole Al Cool
2	0	mati	ure pAPC, immature pAPC, and precursors to pAPC;
	21		exposing the isolated cells to an antigen so that mature pAPC displaying the antigen are
,	22	gen	erated; and
			Express Mail No. <u>EJ455653983US</u> 1009 66

response is inhibited. 110. The method of claim 109, wherein: the step of exposing is performed under conditions selected so that mature pAPC displaying antigen is a produced and a pre-determined set of cytokines, selected from the group consisting of Th1 cytokines and Th2 cytokines, is expressed. 111. The method of claim 109 wherein: the pre-determined T cell response is selected from the group consisting of: a Th1 response and a Th2 response. 12 13. The method of claim 111, wherein: the Th1 or Th2 response is inhibited through induction of an opposing Th2 or Th1
110. The method of claim 109, wherein: the step of exposing is performed under conditions selected so that mature pAPC displaying antigen is a produced and a pre-determined set of cytokines, selected from the group consisting of Th1 cytokines and Th2 cytokines, is expressed. 111. The method of claim 109 wherein: the pre-determined T cell response is selected from the group consisting of: a Th1 response and a Th2 response. 121. The method of claim 111, wherein:
the step of exposing is performed under conditions selected so that mature pAPC displaying antigen is a produced and a pre-determined set of cytokines, selected from the group consisting of Th1 cytokines and Th2 cytokines, is expressed. The method of claim 109 wherein: the pre-determined T cell response is selected from the group consisting of: a Th1 response and a Th2 response. The method of claim 111, wherein:
the step of exposing is performed under conditions selected so that matter pro- displaying antigen is a produced and a pre-determined set of cytokines, selected from the group consisting of Th1 cytokines and Th2 cytokines, is expressed. 111. The method of claim 109 wherein: the pre-determined T cell response is selected from the group consisting of: a Th1 response and a Th2 response. 12 13. The method of claim 111, wherein:
displaying antigen is a produced and a pre-determined set of cytokines, sciented 27 consisting of Th1 cytokines and Th2 cytokines, is expressed. 111. The method of claim 109 wherein: the pre-determined T cell response is selected from the group consisting of: a Th1 response and a Th2 response. 12 13. The method of claim 111, wherein: the Th1 or Th2 response is inhibited through induction of an opposing Th2 or Th1
consisting of Th1 cytokines and Th2 cytokines, is expressed. 111. The method of claim 109 wherein: 12. The method of claim 111, wherein: 13. The method of claim 111, wherein:
consisting of Th1 cytokines and Th2 cytokines, is expressed. 111. The method of claim 109 wherein: 12. The method of claim 111, wherein: 13. The method of claim 111, wherein:
9 111. The method of claim 109 wherein: 10 the pre-determined T cell response is selected from the group consisting of: a Th1 11 response and a Th2 response. 12 13 112. The method of claim 111, wherein:
111. The method of claim 109 wherein: the pre-determined T cell response is selected from the group consisting of: a Th1 response and a Th2 response. The method of claim 111, wherein: Th1 or Th2 response is inhibited through induction of an opposing Th2 or Th1
response and a Th2 response. 12 13 112. The method of claim 111, wherein: 13 the Th1 or Th2 response is inhibited through induction of an opposing Th2 or Th1
response and a Th2 response. 12 13 112. The method of claim 111, wherein: 13 the Th1 or Th2 response is inhibited through induction of an opposing Th2 or Th1
12 13 112. The method of claim 111, wherein: 14. Th1 or Th2 response is inhibited through induction of an opposing Th2 or Th1
13 112. The method of claim 111, wherein:
the Th1 or Th2 response is inhibited through induction of an opposing Th2 or Th2
14
15 response.
16 17 113. The method of claim 109, wherein:
17 113. The method of claim 109, wherem. 18 the step of contacting comprises contacting the antigen-exposed pAPC with T cells in the
the step of contacting complications that a Th2 response is inhibited. 19 presence of one or more Th1 stimulating cytokines, so that a Th2 response is inhibited.
presence of one or more In Stimulating 5,33
20
21 114. The method of claim 109, wherein:

1 2 3	the step of contacting comprises contacting the antigen-exposed pAPC with T cells in the presence of one or more Th1 stimulating cytokines selected from the group consisting of selected from the group consisting of IL-12, IL-18, IL-18, fragments of IL-18, IFN α , and
4 5 6 7 8 9	 115. The method of claim 109, wherein: the step of contacting comprises contacting the antigen-exposed pAPC with T cells in the presence of a Th1 inducing agent, so that the expression of or more Th1 cytokines is induced and a Th2 response is inhibited in the T cells.
10 11 12 13 14 15	is inhibited in the T cells.
2	The method of claim 116, wherein: the microbial extracts are selected from the group consisting of any Staphylococcus

	the step of contacting comprises contacting the mature pAPC displaying antigen with T
1	the step of condemice The stimulating cytokines
2	cells in the presence of one or more Th2 stimulating cytokines
3	
4	119. The method of claim 109, wherein:
5	the step of contacting comprises contacting the mature pAPC displaying antigen with T
6	cells in the presence of one or more cytokines selected from the group consisting of IL-4, so that
7	a Th1 response is inhibited.
8	
9	120. The method of claim 109, wherein:
10	120. The method of claim 1999 the step of contacting comprises contacting the mature pAPC displaying antigen with T
	cells in the presence of one or more Th2 inducing agents.
11	CCII3 M2 man 1
12	121. The method of claim 109, wherein:
13	121. The method of claim 105, was the step of contacting comprises contacting the mature pAPC displaying antigen with T
14	the step of contacting comprises contacting and the induce expression of IL-4 in the
15	cells in the presence of one or more agents selected to induce expression of IL-4 in the
16	responding T cells
17	
18	122. The method of claim 109, wherein:
19	the pAPC are selected from the group consisting of dendritic cells, B cells, and
20	macrophages.
21	l
2:	
	Mail No. F1455653983US

1	t	he pAPC are dendritic cells.
2 3 4 5 6		The method of claim 123, wherein: the step of isolating comprises isolating immature dendritic cells from an individual; and maturing the immature cells <i>in vitro</i> by exposure to one or more cytokines selected from oup consisting of: GM-CSF, IL-3, and IL-4.
7 8 9	125.	The method of claim 123, wherein: the step of maturing is performed concurrently with the step of exposing to antigen.
10 11 12 13	126.	The method of claim 109, wherein: the step of exposing the isolated cells to an antigen comprises exposing the cells to a defended entigen preparation.
14 15 16	12	the doctor of claim 109, wherein:
18 19 20 2)) 11	28. The method of claim 109, wherein: the step of exposing the isolated cells to antigen comprises exposing the cells to a gene encoding the antigen, so that the gene becomes expressed within the cells.
2	22	T1455653983US

1 2 3	129.	The method of claim 125, wherein: the step of exposing further comprises exposing the cells to a factor selected from the consisting of cytokines and inducing agents.
4 5 6 7	130. gene	The method of claim 129, wherein: the factor is a polypeptide and the step of exposing comprises contacting the cells with a encoding the factor.
8 9 10 11	131 a g	. The method of claim 130, wherein: the antigen is a polypeptide and the step of exposing comprises contacting the cells with ene encoding the antigen.
12 13 14 15	13	 The method of claim 131, wherein: the gene encoding the antigen and the gene encoding the factor are coordinately egulated.
16 17 18	1	33. The method of claim 130, wherein: the gene encoding the antigen and the gene encoding the factor are provided on the same nucleic acid molecule.
19 20 2	0	134. The method of claim 132, wherein:

		the gene encoding the antigen and the gene encoding the factor are linked to one another
1		
2	so that	a fusion protein is encoded.
3		
4	135.	The method of claim 132, wherein: the gene encoding the antigen and the gene encoding the factor are provided on separate
5		the gene encoding the antigen and the gene checoung
6	nucle	eic acid molecules.
7		
8	136.	The method of claim 109 wherein:
9		the antigen is provided in association with a targeting agent.
10		
11	137	7. The method of claim 129, wherein:
12		7. The method of claim 129, wherein one or both of the antigen and factor is provided in association with a targeting agent.
13		
14	13	The method of claim 136 or claim 137, wherein: the association with the targeting agent occurs by means of an interaction selected from
15		the association with the targeting agent occurs by meaning the association with the targeting agent occurs by meaning the association with the targeting agent occurs by meaning the association with the targeting agent occurs by meaning the association with the targeting agent occurs by meaning the association with the targeting agent occurs by meaning the association with the targeting agent occurs by meaning the association with the targeting agent occurs by meaning the association with the targeting agent occurs by meaning the association with the targeting agent occurs by meaning the association with the targeting agent occurs by meaning the association with the association and the association are also as a second occurs by the association and the association are also as a second occurs by the association and the association are also as a second occurs by the association and the association are also as a second occurs by the association are also as
16	tł	the association with the day of the day
17	7 h	sydrophobic interactions, and combinations thereof.
1	8	
1	9	139. The method of claim 136 or claim 137, wherein:
	20	139. The method of claim 130 of claim 250, the targeting agent is selected from the group consisting of mannose receptor ligand and
		the Fc receptor ligand.
	22	Toward Mail No. FJ455653983US

1	140.	The method of claim 136 or claim 137, wherein:
1	140.	the targeting agent comprises complement receptor ligand.
2		
4	141.	The method of claim 136 or claim 137, wherein:
5		the targeting agent comprises DEC205.
6		
7	142.	The method of claim 136 or claim 137, wherein:
8		the targeting agent is capable of targeting to particular vesicles within pAPCs.
9		
10	143	The method of claim 136 or claim 137, wherein:
11		the targeting agent comprises at least the Fc portion of an Ig molecule.
12		142 mborgin
13	14	The method of claim 143, wherein: the targeting agent comprises at least the Fc portion of an IgG molecule.
14		the targeting agent comprises at least 2
15		5. The method of claim 109, wherein:
16		5. The method of claim 105, was the step of exposing comprises providing the antigen in an encapsulation device.
17		the step of exposing as a
18		46. The method of claim 129, wherein:
19		one or both of the antigen and factor is encapsulated.
20		
2		147. The method of claim 129, wherein:
2	.4	Express Mail No. EJ455653983US
		April 0 1999 / 3

1		the antigen and factor are provided together as a single composition.
2 3 4	148.	The method of claim 147, wherein: the antigen and factor are provided encapsulated together in a single encapsulation
5	device	2.
6 7 8	149.	The method of claim 145, 146, or claim 148, wherein: the encapsulation device is associated with a targeting agent.
9 10 11	150	The method of claim 149, wherein: the targeting agent is selected from the group consisting of mannose receptor ligand and
12	the	Fc receptor ligand.
13 14 15	15	1. The method of claim 149, wherein: the targeting agent comprises complement receptor ligand.
16 17	15	52. The method of claim 149, wherein:
18		the targeting agent comprises DEC205.
19		
20 21		53. The method of claim 149, wherein: the targeting agent is capable of targeting to intracellular vesicles within pAPCs.
2		T Moil No. F1455653983US

1	154.	7	The method of claim 149, wherein:
2		1	the targeting agent comprises at least the Fc portion of an Ig molecule.
3			
4	155.		The method of claim 149, wherein:
5			the targeting agent comprises at least the Fc portion of an IgG molecule.
6			
7	156	5.	The method of claim 129, wherein:
8			the step of exposing comprises providing antigen and factor that are associated with one
9	ano	oth	er by means of an interaction selected from the group consisting of: covalent bonds,
10	hy	dro	gen bonds, van der Waals interactions, hydrophobic interactions, and combinations thereof.
11			
12	15	57.	The method of claim 109, wherein:
13			the step of exposing the antigen comprises exposing the cells to a modified antigen.
14			
15		58.	The method of claim 149, wherein:
16			the antigen comprises an autoantigen; and
13			the pre-determined set of cytokines comprises Th2 cytokines.
1			
		159	9. The method of claim 149, wherein:
	20		the pre-determined set of cytokines comprises IL-4.
	21		-
2	22		TI4556530831IS

1	160.	A method of treating allergy, the method comprising steps of:
2		identifying an individual who is allergic to an antigen;
3		providing a composition of pAPC displaying the antigen; and
4		contacting the composition with T cells of the individual under conditions that inhibit a
5	Th2 re	esponse to the antigen.
6		
7	161.	The method of claim 160, wherein:
8	102	the mature pAPC are selected for their expression of Th1 cytokines.
9		
10	162.	The method of claim 160, wherein:
	102.	the pAPC are selected from the group consisting of dendritic cells, B cells, and
11	mac	rophages.
12	mac	Topings
13	163	. The method of claim 161, wherein:
14	103	the pAPC are dendritic cells.
15		the price of the second
16	• •	4. The method of claim 160, wherein:
17	164	the step of providing comprises:
18		isolating from an individual one or more cells selected from the group consisting
19		mature pAPC, immature pAPC, and precursors to pAPC; and
20	of	exposing the isolated cells to the antigen.
21		exposing the isolated cells to the many
22		N. 121. E14556539831IS

1	165.	The method of claim 164, wherein:
2		the step of exposing the isolated cells to the antigen further comprises exposing the
3	isolate	ed cells to a factor selected from the group consisting of cytokines and inducing agents.
4		
5	166.	The method of claim 165, wherein:
6		the factor comprises an inducing agent that induces expression of one or more Th1
7	stimu	llating cytokines in the pAPC.
8		
9	167.	The method of claim 165 wherein:
10		the antigen and factor are provided together as part of a single composition.
11		
12	168	. The method of claim 165, wherein:
13		one or both of the antigen and factor is associated with a targeting agent.
14		
15	169	The method of claim 164, wherein:
16		the antigen is associated with a targeting agent.
17		
18	17	0. The method of claim 167, wherein:
19		the antigen and factor are encapsulated together in an encapsulation device.
20		
21	17	71. The method of claim 164, wherein
22		the antigen is encapsulated.

1	172.	The method of claim 165, wherein:
2		one or both of the antigen and factor is encapsulated.
3		
4	173.	The method of claim 165, wherein:
5		the antigen and factor are both encapsulated.
6		
7	174.	The method of claim 173, wherein:
8		the encapsulation device is associated with a targeting agent.
9		
10	175	The method of claim 164, wherein:
11		the step of exposing the isolated cells to antigen comprises exposing the cells to a crude
12	pre	paration of antigen.
13		
14	170	6. The method of claim 164, wherein:
15		the step of exposing the isolated cells to an antigen comprises exposing the cells
16	su	bstantially pure antigen.
17		
18	1	77. The method of claim 164, wherein:
19	1	the antigen is a polypeptide antigen; and
20)	the antigent is a possperation that the step of exposing the isolated cells to antigen comprises exposing the cells to a gene
2	1 6	encoding the antigen, so that the gene becomes expressed within the cells.
2		

1 2	178.	The method of claim 164, wherein: the factor is a polypeptide and the step of exposing comprises exposing the cells to a gene
3	enco	ding the factor.
4		
5	179.	
6		the antigen is a polypeptide antigen; and
7		the antigen is a posspection of the antigen comprises exposing the cells to a gene the step of exposing the isolated cells to antigen comprises exposing the cells to a gene
8	ence	oding the antigen, so that the gene becomes expressed within the cells.
9		
10	180). The method of claim 179, wherein:
11		the antigen gene and the factor gene are coordinately regulated.
12		
13	18	31. The method of claim 179, wherein:
14		the antigen gene and the factor gene are provided on the same nucleic acid molecule.
15		
16	1	82. The method of claim 181, wherein:
17		the antigen gene and the factor gene are linked to one another so that a single fusion
18	ŗ	protein is encoded.
19)	
20)	183. The method of claim 179, wherein:
2		the antigen gene and the factor gene are provided on separate nucleic acid molecules.
	2	
_		Nail No. 51455653983US

1	184.	The method of any one of claims 168, 169, or 174, wherein:
2		the association with the targeting agent occurs through an interaction selected from the
3	group	consisting of covalent bonds, hydrogen bonds, van der Waals interactions, hydrophobic
4		ctions, and combinations thereof.
5		
6	185.	The method of any one of claims 168, 169, or 174, wherein:
7	102.	the targeting agent is selected from the group consisting of mannose receptor ligand and
8	the F	c receptor ligand.
9		
10	186.	The method of any one of claims 168, 169, or 174, wherein:
11		the targeting agent comprises complement receptor ligand.
12		
13	187	The method of any one of claims 168, 169, or 174, wherein:
14		the targeting agent comprises DEC205.
15		
16	18	8. The method of any one of claims 168, 169, or 174, wherein:
17		the targeting agent is capable of targeting to intracellular vesicles within pAPCs.
18		
19	18	39. The method of any one of claims 168, 169, or 174, wherein:
20		the targeting agent comprises at least the Fc portion of an Ig molecule.
21		
21	1	90. The method of any one of claims 168, 169, or 174, wherein:
22	. •	Fueress Mail No. F1455653983 <u>US</u>

1		the targeting agent comprises at least the Fc portion of an IgG molecule.
2		
3	191.	The method of claim 175, wherein: the step of exposing comprises providing antigen and factor that are associated with one the step of exposing comprises providing antigen and factor that are associated with one that the group consisting of: covalent bonds,
5	anothe	er by means of an interaction selected from the group consisting of: covalent bonds,
6	hydro	gen bonds, van der Waals interactions, hydrophobic interactions, and combinations thereof.
7 8 9	192.	The method of claim 164, wherein: the step of exposing the antigen comprises exposing the cells to a modified antigen.
101112	193.	The method of claim 192, wherein: the modified antigen is substantially identical to a naturally-occurring antigen that ains at least one IgE binding site except that the modified antigen lacks at least one of the
13 14		
15 16 17 18 19 20	194	1. Starting on autoimmune disorder, the method comprising steps of:
21 22	Th	1 response to the antigen.

1	195.	The method of claim 194, wherein:	
2		the step of identifying comprises identifying an individual who has previously mounted a	
3	Th1 re	sponse to the antigen.	
4			
5	196.	The method of claim 194, wherein:	
6		the pAPC are selected for their expression of Th2 stimulating cytokines.	
7			
8	197.	The method of claim 194, wherein:	
9		the pAPC are selected from the group consisting of dendritic cells, B cells, and	
10	macrophages.		
11			
12	198.	The method of claim 194, wherein:	
13		the pAPC are B cells.	
14			
15	199.	The method of claim 194, wherein:	
16		the step of providing comprises:	
17		isolating from an individual one or more cells selected from the group consisting	
18	of n	nature pAPC, immature pAPC, and precursors to pAPC; and	
19		exposing the isolated cells to the antigen.	
20			
21	200	The method of claim 199, wherein:	

		the step of exposing the isolated cells to the antigen further comprises exposing the
1		
2	isolated	d cells to a factor selected from the group consisting of cytokines and inducing agents.
3		
4	201.	The method of claim 200, wherein:
5		the factor comprises an inducing agent that induces expression of one or more Th2
6	cytoki	nes.
7		
8	202.	The method of claim 200, wherein:
9		the antigen and factor are provided together as part of a single composition.
10		
11	203.	The method of claim 200, wherein:
12		one or both of the antigen and factor is associated with a targeting agent.
13		
14	204.	The method of claim 199, wherein:
15		the antigen is associated with a targeting agent.
16		
17	205.	The method of claim 203, wherein:
18		the antigen and factor are encapsulated together in an encapsulation device.
19		
20	206.	The method of claim 199, wherein
21		the antigen is encapsulated.
22		

1	207.	The method of claim 200, wherein:
2		the antigen and factor are both encapsulated.
3		
4	208.	The method of claim 205, 206, or 207 wherein:
5		the encapsulation device is associated with a targeting agent.
6		
7	209.	The method of claim 200, wherein:
8		the step of exposing the isolated cells to antigen comprises exposing the cells to a crude
9	prepa	aration of antigen.
10	•	
11	210.	The method of claim 200, wherein:
12		the step of exposing the isolated cells to an antigen comprises exposing the cells
13	sub	stantially pure antigen.
14		
15	211	. The method of claim 199, wherein:
16		the antigen is a polypeptide antigen; and
17		the step of exposing the isolated cells to antigen comprises exposing the cells to a gene
18	ene	coding the antigen, so that the gene becomes expressed within the cells.
19		
20	21	2. The method of claim 200, wherein:
		the factor is a polypeptide and the step of exposing comprises exposing the cells to a gene
21		acoding the factor.
22	, ei	Express Mail No. <u>EJ455653983US</u>
		Express than 1.5

1	213.	The method of claim 212, wherein:
2		the antigen is a polypeptide antigen; and
3		the step of exposing the isolated cells to antigen comprises exposing the cells to a gene
4	encodi	ng the antigen, so that the gene becomes expressed within the cells.
5		
6	214.	The method of claim 213, wherein:
7		the antigen gene and the factor gene are coordinately regulated.
8		
9	215.	The method of claim 213, wherein:
10		the antigen gene and the factor gene are provided on the same nucleic acid molecule.
11		
12	216.	The method of claim 215, wherein:
13		the antigen gene and the factor gene are linked to one another so that a single fusion
14	prote	ein is encoded.
15		
16	217.	The method of claim 213, wherein:
17		the antigen gene and the factor gene are provided on separate nucleic acid molecules.
18		
19	218	The method of any one of claims 203, 204, or 208, wherein:
20		the association with the targeting agent occurs through an interaction selected from the
21	gro	up consisting of covalent bonds, hydrogen bonds, van der Waals interactions, hydrophobic
22	inte	eractions, and combinations thereof.

1	219.	The method of any one of claims 203, 204, or 208, wherein:
2		the targeting agent is selected from the group consisting of mannose receptor ligand and
3	the Fc	receptor ligand.
4		
5	220.	The method of any one of claims 203, 204, or 208, wherein:
6		the targeting agent comprises complement receptor ligand.
7		
8	221.	The method of any one of claims 203, 204, or 208, wherein:
9		the targeting agent is capable of targeting to intracellular vesicles within pAPCs.
10		
11	222.	The method of any one of claims 203, 204, or 208, wherein:
12		the targeting agent comprises at least the Fc portion of an Ig molecule.
13		
14	223.	The method of any one of claims 203, 204, or 208, wherein:
15		the targeting agent comprises at least the Fc portion of an IgG molecule.
16		
17	224.	
18		the step of exposing comprises providing antigen and factor that are associated with one
19	ano	ther by means of an interaction selected from the group consisting of covalent bonds, van der
20	Wa	als interactions, hydrophobic interactions, and combinations thereof.
21		
22	225	The method of claim 199, wherein:

1		the step of exposing the antigen comprises exposing the cells to a modified antigen.
2	226	The method of claim 225, wherein:
3	226.	the modified antigen is substantially identical to a naturally-occurring antigen that
4		
5	contai	ns at least one IgE binding site except that the modified antigen lacks at least one of the
6	IgE bi	nding sites.
7		
8	227.	A composition for modulating an immune system response to an antigen in an individual
9	comp	rising:
10		an antigen; and
11		at least one factor selected from the group consisting of cytokines and inducing
12	agen	ts.
13		·
14	228.	The composition of claim 227, wherein:
15		the factor comprises a Th1 stimulating cytokine.
16		
17	229	
18		the factor is selected from the group consisting of IL-12, IL-2, IL-18, IL-1B, fragments
19	of I	$_{-1}$ B, IFN α , and IFN γ .
20		
21	230	. The composition of claim 227, wherein:
22		the factor comprises a Th2 stimulating cytokine.
		Former Mail No. FI455653983US

1	231.	The composition of claim 227, wherein:
2		the factor comprises IL-4.
3		
4	232.	The composition of claim 227, wherein:
5		the factor comprises a Th1 inducing agent.
6		
7	233.	The composition of claim 227, wherein:
8		the factor is selected from the group consisting of LPS, CD40, CD40 ligand, BCGs,
9	oligor	nucleotides containing CpG motifs, TNF α , and microbial extracts.
10		
11	234.	The composition of claim 233, wherein:
12		the microbial extracts are selected from the group consisting of any Staphylococcus
13	aurei	as preparation, heat killed Listeria, and modified cholera toxin.
14		
15	235.	The composition of claim 227, wherein:
16		the factor comprises a Th2 inducing agent.
17		
18	236.	The composition of claim 227, wherein:
19		the factor comprises an agent that induces IL-4 expression.
20		
21	237.	The composition of claim 227, wherein:
22		the antigen comprises a crude antigen preparation.

1	238.	The composition of claim 227, wherein:
2		the antigen comprises a substantially pure antigen
3		
4	239.	The composition of claim 227, further comprising:
5		an encapsulation device surrounding the antigen and factor.
6		
7	240.	The composition of claim 227 or claim 228, further comprising:
8		a targeting agent.
9		
10	241.	The composition of claim 240, wherein:
11		the targeting agent is associated with the composition through a covalent or a non-
12	coval	ent interaction.
13		
14	242.	The composition of claim 240, wherein:
15		the targeting agent is selected from the group consisting of mannose receptor ligand and
16	the F	c receptor ligand.
17		
18	243.	-
19		the targeting agent comprises complement receptor ligand.
20		
21	244.	
22		the targeting agent comprises DEC205.

1	245.	The composition of claim 239, wherein:
2		the targeting agent is capable of targeting to intracellular vesicles within pAPCs.
3		
4	246.	The composition of claim 239, wherein:
5		the targeting agent comprises at least the Fc portion of an Ig molecule.
6		
7	247.	The composition of claim 239, wherein:
8		the targeting agent comprises at least the Fc portion of an IgG molecule.
9		
10	248.	The composition of claim 227, wherein:
11		the antigen and factor are covalently linked to one another.
12		
13	249.	The composition of claim 227, wherein:
14		the antigen and factor that are associated with one another by means of an interaction
15	selec	ted from the group consisting of: hydrogen bonds, van der Waals interaction, hydrophobic
16	intera	action, and combinations thereof.
17		
18	250.	The composition of claim 227, wherein:
19		the antigen comprises a modified antigen.
20		
21	251.	The composition of claim 227, which composition is formulated for oral administration.
22		

1	252.	The composition of claim 227, which composition is formulated for inhalation.
2	202.	•
3	253.	The composition of claim 227, which composition is formulated for injection.
4		
5	254.	A composition for modulating an immune system response to an antigen in an individual
6	compr	ising:
7		one or more pAPC displaying an antigen and expressing a predetermined collection of
8	cytoki	nes, selected from the group consisting of Th1 cytokines and Th2 cytokines; and
9		at least one factor selected from the group consisting of cytokines and inducing agents.
10		
11	255.	The composition of claim 254, wherein:
12		the pAPC are selected form the group consisting of dendritic cells, B cells, and
13	macro	ophages.
14		
15	256.	The composition of claim 255, wherein:
16		the pAPC are dendritic cells.
17		
18	257.	The composition of claim 255, wherein:
19		the dendritic cells are prepared by a process comprising steps of:
20		isolating immature dendritic cells from an individual; and
21		maturing the isolated cells in vitro by exposure to one or more cytokines selected
22	from	the group consisting of: GM-CSF, IL-3, and IL-4.

1	258.	The composition of 257, wherein:
2		the maturing is performed in the presence of the antigen.
3		
4	259.	The composition of claim 254, wherein:
5		the factor comprises a Th1 stimulating cytokine.
6		
7	260.	The composition of claim 254, wherein:
8		the factor is selected from the group consisting of IL-12, IL-2, IL-18, IL-1B, fragments
9	of IL-	1ß, IFN α , and IFN γ
10		
11	261.	The composition of claim 254, wherein:
12		the factor comprises a Th1 inducing agent.
13		
14	262.	
15		the factor is selected from the group consisting of LPS, CD40, CD40 ligand, BCGs,
16	oligo	onucleotides containing CpG motifs, TNFα, and microbial extracts.
17		
18	263	
19		the microbial extracts are selected from the group consisting of any Staphylococcus
20	aur	eus preparation, heat killed Listeria, and modified cholera toxin.
21		
22	264	The composition of claim 254, wherein:
	Apr	Express Mail No. <u>EJ455653983US</u>

1		the factor comprises a Th2 stimulating cytokine.
2		
3	265.	The composition of claim 254, wherein:
4		the factor comprises IL-4.
5	•	
6	266.	The composition of claim 254, wherein:
7		the factor comprises a Th2 inducing agent.
8		
9	267.	The composition of claim 254, wherein:
10		the factor comprises an agent that induces IL-4 expression.
11		
12	268.	The composition of claim 254, wherein:
13		the factor comprises an agent that inhibits IL-12 expression.
14		
15	269.	The composition of claim 258, wherein:
16		the antigen comprises a crude antigen preparation.
17		
18	270.	The composition of claim 254, wherein:
19		the antigen comprises a substantially pure antigen.
20		
21	271.	The composition of claim 254, wherein:
22		the antigen comprises a modified antigen.

1	272.	A composition comprising:
2		a gene encoding an antigen; and
3		a gene encoding at least one factor selected from the group consisting of cytokines and
4	induci	ing agents.
5		
6	273.	The composition of claim 272, wherein:
7		the antigen gene and the factor gene are coordinately regulated.
8		
9	274.	The composition of claim 272, wherein:
10		the antigen gene and the factor gene are on the same nucleic acid molecule.
11		
12	275.	The composition of claim 272, wherein:
13		the antigen gene and the factor gene are linked together so that a single polypeptide is
14	encod	led.
15		
16	276.	The composition of claim 272, wherein:
17		the antigen gene and the factor gene are provided on separate nucleic acid molecules.
18		
19	277.	The composition of claim 272, further comprising an encapsulation device surrounding
20	the go	enes.
21		
22	278.	The composition of claim 272 or claim 277, further comprising:

1		a targeting agent selected for its ability to localize the composition in the vicinity of
2	pAPC.	
3		
4	279.	The composition of claim 272, which composition is formulated for oral administration.
5		
6	280.	The composition of claim 272, which composition is formulated for inhalation.
7		
8	281.	The composition of claim 272, which composition is formulated for injection.